Face your challenge, Be smart



THEORETICAL EXAMINATION

JULY 20, 2013 MOSCOW, RUSSIA

General Directions

- Write down your name and code number on each page.
- You have 5 h to fulfill the tasks. Failure to stop after the STOP command may result in zero points for the current task.
- Write down answers and calculations within the designated boxes. Give your work where required.
- Use only the pen and calculator provided.
- If you need draft paper use the back side of the paper. It will not be marked.
- There are **38** pages in the booklet including the answer boxes, Cover Sheet and Periodic Table.
- The official English version is available on demand for clarification only.
- Need to go to the restroom raise your hand. You will be guided there.
- After the STOP signal put your booklet in the envelope (don't seal), leave at your table. Do not leave the room without permission.
- You have additional 15 minutes to read the whole set.

Avogadro's constant	$N_{\rm A} = 6.0221 \times 10^{23} {\rm mol}^{-1}$
Universal gas constant	$R = 8.3145 \text{ J} \cdot \text{K}^{-1} \cdot \text{mol}^{-1}$
Speed of light	$c = 2.9979 \times 10^8 \text{ m} \cdot \text{s}^{-1}$
Planck's constant	$h = 6.6261 \times 10^{-34} \mathrm{J} \cdot \mathrm{s}$
Faraday constant	$F = 96485 \text{ C} \cdot \text{mol}^{-1}$
Gravity of Earth	$g = 9.81 \text{ m} \cdot \text{s}^{-2}$
Standard pressure	$p^{\circ} = 1$ bar $= 10^5$ Pa $= 750$ mmHg
Atmospheric pressure	$1 \text{ atm} = 1.013 \times 10^5 \text{ Pa} = 760 \text{ mmHg}$
Zero of the Celsius scale	273.15 K

Physical Constants, Units, Formulas and Equations

1 nanometer (nm) = 10^{-9} m

1 Da = 1 atomic mass unit

Energy of a light quantum with wavelength λ $E = hc / \lambda$ Energy of one mole of photons $E_{\rm m} = hcN_{\rm A} / \lambda$ G = H - TSGibbs energy $K = \exp\left(-\frac{\Delta G^{\circ}}{RT}\right)$ Relation between equilibrium constant and standard Gibbs energy Relation between standard Gibbs energy and $\Delta G^{\circ} = -nFE^{\circ}$ standard emf $\frac{dp}{dT} = \frac{\Delta H}{T\Delta V}$ Clapeyron equation for phase transitions $\ln\frac{p_2}{p_1} = \frac{\Delta H}{R} \left(\frac{1}{T_1} - \frac{1}{T_2}\right)$ Integrated Clausius-Clapeyron equation for phase transitions involving vapor $\Delta G = \Delta G^{\circ} + RT \ln \frac{a_{\rm prod}}{a_{\rm reag}},$ Dependence of Gibbs energy of reaction on concentration or pressure a = c / (1 mol/L) for the substances in solution, a = p / (1 bar) for gases $V = \frac{4}{3}\pi R^3$ Volume of a sphere of radius *R* $S = 4\pi R^2$ Surface area of a sphere of radius *R* $p = \rho g h$ Hydrostatic pressure

1 electron volt (eV) = $1.6022 \cdot 10^{-19} \text{ J} = 96485 \text{ J} \cdot \text{mol}^{-1}$

Problem 1. Clathrate gun (8 points)

Question	1	2	3	4	5	6	Total
Marks	2	1	3	5	6	2	19

The only gun that is able to kill all living people in one shot

On the floors of oceans and seas there are vast reserves of methane in the form of clathrate compounds called methane hydrates. These reserves can be mined and serve as a source of energy or raw materials for organic synthesis. However, scientists are seriously worried about the possibility of spontaneous decomposition of hydrates caused by the raising ocean temperature. It is believed that if a sufficient amount of methane is released into the atmosphere, the oceans will warm up quicker due to the greenhouse effect, further accelerating the decomposition of clathrates. Due to the



explosion of the resulting methane-air mixture and/or changes in the composition of the atmosphere, all living creatures may become extinct. This apocalyptic scenario is called a clathrate gun.

Upon decomposition of 1.00 g of a methane hydrate with a fixed composition at 25 °C and atmospheric (101.3 kPa) pressure, 205 mL of methane is released.

1. Determine *n* (not necessarily integer) in the formula of methane hydrate, $CH_4 \cdot nH_2O$.

Calculations:

Answer:

Real methane hydrate has a non-stoichiometric composition close to $CH_4 \cdot 6H_2O$. At atmospheric pressure, methane hydrate decomposes at -81 °C. However, under high pressures (e.g. on the ocean floor) it is stable at much higher temperatures. Decomposition of methane hydrate produces gaseous methane and solid or liquid water depending on temperature.

2. Write down the equation of decomposition of 1 mole of $CH_4 \cdot 6H_2O$ producing solid water (ice) $H_2O(s)$.

The enthalpy of this process equals $17.47 \text{ kJ} \cdot \text{mol}^{-1}$. Assume that the enthalpies do not depend on temperature and pressure, the volume change upon decomposition of hydrate is equal to the volume of released methane, and methane is an ideal gas.

3. At what external pressure does decomposition of methane hydrate into methane and ice take place at -5 °C?

Calculations:	
Answer:	

4. What is the minimum possible depth of pure liquid water at which methane hydrates can be stable?

To answer this question, you should first deduce at which minimum temperature methane hydrate can coexist with liquid water. Choose the correct answer.

□ 272.9 К	273.15 К	273.4 К

Calculations:

Answer:

Large methane hydrate stocks on the floor of Baikal lake, the largest freshwater lake in Russia and in the world, have been discovered in July 2009 by the crew of a deep-submergence vehicle «Mir-2». During the ascent from the depth of 1400 m methane hydrate samples started to decompose at the depth of 372 m.

5. Determine the temperature in Baikal lake at the depth of 372 m. The enthalpy of fusion of ice is $6.01 \text{ kJ} \cdot \text{mol}^{-1}$.

Calculations:

Answer:

Total amount of methane in hydrates on Earth is no less than $5 \cdot 10^{11}$ tons.

6. By how many degrees would the Earth atmosphere heat up, if such amount of methane is burned by reacting with atmospheric oxygen? The enthalpy of combustion of methane is - 889 kJ·mol⁻¹, the total heat capacity of the Earth's atmosphere is about $4 \cdot 10^{21}$ J·K⁻¹.

Calculations:

Answer:

Problem 2. Break down photosynthesis – the Hill reaction (7 points)

Question	1	2	3		2	1	5 6		Total	
Question	1	Z	a	b	с	a	b	5	0	Total
Points	1	2	2	2	3.5	1	2	3	2.5	19

In the history of photosynthesis research, there were some breakthrough experiments which added much to our knowledge of this very complex process. One of such experiments was performed in 1930s by an English biochemist Robert Hill. In this problem, we consider some of his data together with the data of more recent experiments.

1. In plants, under illumination, carbon dioxide is reduced to carbohydrates (denote as $\{CH_2O\}$) and oxygen is produced. Write the overall equation of photosynthesis in plants.

Much of the photosynthesis takes place in chloroplasts – organelles found in plant cells and containing chlorophyll – the light-absorbing substance. Hill isolated chloroplasts from the cells by grinding the leaves in the sucrose solutions. The cell-free chloroplasts did not produce oxygen under illumination even in the presence of CO_2 . However, upon adding potassium ferrioxalate $K_3[Fe(C_2O_4)_3]$ (with the excess of potassium oxalate) to the chloroplast suspension Hill observed oxygen liberation under illumination even without CO_2 .

2. Hill's experiment enabled to determine the source of oxygen during photosynthesis. Write the formulas of the oxidant and the reducing agent in the photosynthesis inside the plant cells and in the cell-free chloroplasts (the Hill reaction).

Natural	Natural photosynthesis		reaction
Oxidant	Reducing agent	Oxidant	Reducing agent

Hill measured the amount of evolved oxygen using muscle haemoglobin (Hill denoted it Hb) which binds all molecular oxygen in a 1:1 ratio to form HbO₂. The initial concentration of Hb was $0.6 \cdot 10^{-4}$ M. Kinetic curves corresponding to different ferrioxalate concentrations are shown in the figure (the upper curve corresponds to $2.0 \cdot 10^{-4}$ M).



The fraction of bound haemoglobin HbO_2 (with respect to the initial amount of Hb) as function of time. Crosses denote the end of reaction

(Figure 2a from the original Hill's paper: R. Hill. Oxygen produced by isolated chloroplasts. – Proc. R. Soc. B, 1939, v. 127, pp. 192-210)

3. a. From the figure, estimate the Fe / O_2 mole ratio at the end of reaction. Do not take into account the iron from Hb.

b. Write the equation of Hill reaction assuming that it proceeds with a high yield.

c. Using the table of standard electrode potentials, determine the Gibbs energy of the Hill reaction at T = 298 K, oxygen pressure 1 mmHg, pH = 8 and standard concentrations of other species. Is this reaction spontaneous at such conditions?

Half-reaction	E°, V
$O_2 + 4H^+ + 4e \rightarrow 2H_2O$	+1.23
$\mathrm{CO}_2 + 4\mathrm{H}^+ + 8e \rightarrow \{\mathrm{CH}_2\mathrm{O}\} + \mathrm{H}_2\mathrm{O}$	-0.01
$\mathrm{Fe}^{3+} + e \rightarrow \mathrm{Fe}^{2+}$	+0.77
$\mathrm{Fe}^{3+} + 3e \rightarrow \mathrm{Fe}^{0}$	-0.04
$[\operatorname{Fe}(\operatorname{C}_{2}\operatorname{O}_{4})_{3}]^{3-} + e \to [\operatorname{Fe}(\operatorname{C}_{2}\operatorname{O}_{4})_{3}]^{4-}$	+0.05
$[\text{Fe}(\text{C}_2\text{O}_4)_3]^{4-} + 2e \rightarrow \text{Fe} + 3\text{C}_2\text{O}_4^{2-}$	-0.59

a. Calculations

 $n(Fe) / n(O_2) =$

b.			
Reaction equation:			
c. Calculations			
$\Delta G =$			
The reaction is			
	spontaneous	not spontaneous	

Now, the name "Hill reaction" denotes photochemical oxidation of water by any oxidant other than carbon dioxide which is sensitized by plant cells or isolated chloroplasts.

In another experiment (1952), quinone in an acid solution was used as an oxidant in the Hill reaction initiated by light flashes in the *Chlorella* algae. Experimental data are shown in the figure. The volume of oxygen (in mm³, at temperature 10 °C and pressure 740 mmHg) per one gram of chlorophyll per one flash was determined as a function of light intensity for natural photosynthesis and for isolated chloroplasts. It was found that the maximum yield of oxygen is the same for natural photosynthesis and the Hill reaction.



(Figure 1 from: *H. Ehrmantraut, E. Rabinovitch. Kinetics of Hill reaction.* – *Archives of Biochemistry and Biophysics, 1952, v. 38, pp. 67-84*)

4 a. Determine the reaction order of a photochemical Hill reaction with respect to light intensity at low and high intensity. For each case choose one of three values:

Reaction order:			
Low intensity		High intensity	7
	2		2

b. How many chlorophyll molecules participate in the formation of one oxygen molecule in the saturation limit of the Hill reaction? (The molecular mass of chlorophyll is about 900 Da).

Calculations:	

 $n(Chl) / n(O_2) =$

The quantum requirement of the light redox reactions is defined as the average number of light photons (not necessarily integer) needed for the transfer of one electron from a reducing agent to an oxidant. The isolated chloroplasts were irradiated during 2 hours by a monochromatic light (wavelength 672 nm) with the energy input 0.503 mJ/s, and the total volume of oxygen formed was 47.6 mm³ (under the same conditions as in question 4).

5. Calculate the quantum requirement for the Hill reaction.

Calculations:

Quantum requirement:

6. Try to make conclusions from the above experiments (questions 2-5). For each of the following statements <u>choose either "Yes" or "No"</u>.

	Yes	No
In natural photosynthesis, water oxidation and CO ₂		
reduction are separated in space.		
In chloroplasts, O_2 is produced from CO_2 .		
Oxidation of water in chloroplasts requires light		
illumination.		
Most of chlorophylls in chloroplasts participate directly		
in the photochemical O_2 production.		
In isolated chloroplasts, every absorbed photon causes		
transfer of one electron.		

Problem 3. Meerwein-Schmidt-Ponndorf-Verley reaction (8 points)

Question	1		2	3	4	Total
	а	b				
Marks	7	3	8.5	6	8	32.5

Meerwein-Schmidt-Ponndorf-Verley (MSPV) reaction is a useful tool for reduction of carbonyl compounds to alcohols. The reaction is the reduction of carbonyl compounds by low molecular weight alcohols in the presence of alkoxides of aluminium or other metals:



The mechanism of the reaction includes coordination of carbonyl compound by aluminium alkoxide, hydride transfer in the inner sphere of the complex and subsequent transalkoxylation. It can be schematically represented as follows (transalkoxylation is shown as a one-step process for brevity):



The reaction is reversible and shifting the equilibrium to the desired product requires some excess of the reductant. In some cases (e.g. in the case of reduction of aromatic aldehydes and ketones) the equilibrium constant is so large that the reverse reaction can be neglected.

The	table	below	contains	standard	entropies	and	standard	enthalpies	of	formation	of	liquid
sub	stances	s at 298	K. The b	oiling poi	ints of the	subs	tances at	1 bar are als	50 g	given.		

Substance	$\Delta_f H^{o}_{298}, \text{kJ/mol}$	S^{o}_{298} , J/(mol·K)	$t_{\rm vap}$, °C
Acetone	-248.4	200.4	56
Isopropanol	-318.1	180.6	82
Cyclohexanone	-271.2	229.0	156
Cyclohexanol	-348.2	203.4	161

1a. Calculate the minimum isopropanol:cyclohexanone mass ratio which is required to reach a 99% yield of reaction at 298 K. Assume that a) the reaction mixture eventually gets at equilibrium and b) no products are initially present.

Calculations:

Answer: $m(C_3H_8O) : m(C_6H_{10}O) =$

1b. Choose the appropriate way(s) to increase the cyclohexanol yield.

Warning: erroneously ticked boxes will result in penalty points

Increase the temperature up to 60°C, ev	aporating (distilling)
the acetone	
Add some ethanol to the reaction mixture	
Add some ethanal to the reaction mixture	

2. Often the rate-limiting step in the MSPV reaction is the hydride transfer or the alcoholysis of the alkoxide after hydride transfer. For these two cases, using the above mechanism (2), derive an expression for the rate of reaction as a function of current concentrations of a carbonyl compound, isopropanol and a catalyst. In both cases determine the rate orders in the reactants and the catalyst. Assume that all reaction steps before the limiting step are fast and reversible. Use equilibrium approximation, if necessary. For brevity use the following

notation: A for carbonyl compound, B for isopropanol, C for catalyst. Denote intermediates as you wish.

Rate-limiting step is the hydride transfer

Derivation:

Answer
Order in carbonyl compound:
Order in isopropanol:
Order in the catalyst:

Rate-limiting step is the transalkoxylation of the alcoholate by isopropanol

Derivation:

r =

r =	
A	
Answer	
Order in carbonyl compound:	
Order in isopropanol:	
Order in the catalyst:	

MSPV reaction can be used to obtain chiral alcohols, if the chiral catalyst is employed. For instance, Campbell et al. used the catalyst based on the chiral 2,2'-dihydroxy-1,1'-binaphtyl (BINOL), which is synthesized *in situ* from binaphtol and trimethylaluminium:



(3)

The chirality of BINOL is due to the sterically hindered rotation around the C-C bond. Though perfectly stable at room temperature, BINOL may racemize when heated.

3. Which of the phenols below can form stable (at room temperature) enantiomers so that they can be used in the same fashion to produce a chiral catalyst? *Warning: erroneously ticked boxes will result in penalty points*

Substance	Can be used	Substance	Can be used
OH OH OCH ₃ OCH ₃		ОН	



4. *Enantiomeric excess, ee*, is used to characterize the enantiomeric purity of the substance. This quantity equals ratio of the difference of concentrations of enantiomers R and S to their sum:

 $ee = \frac{[R] - [S]}{[R] + [S]}$

Enantiomeric excess of the pure *R* isomer is unity, *ee* of the racemic mixture is zero.

When using the enantiomerically pure (BINOL)Al(OiPr) as a catalyst for reduction of α bromoacetophenone, the *ee* of the product equals 81%. What is the *ee* of the product if the catalyst *ee* equals 50%? Provide your calculation with an illustration or derivation of the final formula.

Derivation:

ee =

Problem 4. A simple inorganic experiment (6 points)

Question	1	2	3	Total
Marks	5	12	7	24

Compound **A** which contains metal **X** is a colorless crystalline solid and highly soluble in water. It is used as a reagent in analysis and gives in alkali media a binary compound **B** containing 6.9 % (mass) of oxygen. Under heating **A** decomposes with a mass loss of 36.5%.

I. Determine th	he metal \mathbf{X} and compound	$\mathbf{A}, \mathbf{B}.$	
Your work:			
X =	A =	B =	

2. Upon adding some amount of sodium thiosulphate to the solution of **A** the color immediately becomes red, then changes to reddish-brown, and after some minutes a darkbrown precipitate **C** forms (reaction 1). The solution over it is colorless. Being heated on air at 600°C, **C** gives a grey powder **X** (reaction 2), so as 0.90 g of residue can be obtained from 1.10 g of **C**. A gas evolved by heating **C** in vacuum (reaction 3) can be absorbed by calcium hydroxide suspension (reaction 4). Being stored for a long time under saturated solution of barium perchlorate in 0.1 M HClO₄, the color of the precipitate becomes lighter, while the use of magnesium perchlorate doesn't give such effect. What is **C**? Write the equations of the reactions (1 - 4).

Your work:

C = _____

Reaction equations:

3. The compound **C** being stored under the mother liquor (containing an excess of **A**) its color changes to yellow due to the transformation into **D**. If barium ions are added to the suspension of **C** in the mother liquor, a mixture of **D** and of a white precipitate forms. Propose the formula of **D**, taking into account that it contains 77.5% (mass) of **X**. Give the equation of **D** formation.

Your work:

D = _____

Reaction equation:

Problem 5. Simple estimates of graphene properties (7 points)

Question	1		2	3	Total
	а	b			
Marks	2	2.5	4	5.5	14

Graphene is a two-dimensional, one atom thick carbon material (Fig.1 a). Many layers of graphene stack together to form graphite (Fig. 1b).



Fig. 1. (a) The structure of graphene. Spheres are carbon atoms. They are arranged in hexagons. The area of one carbon hexagon is $5.16 \cdot 10^{-20} \text{ m}^2$ (b) Crystal lattice of graphite. Three graphene layers are shown

Such atomic structure was long considered to be unstable. However, in 2004 Andrey Geim and Konstantin Novoselov have reported production of the first samples of this unusual material. This groundbreaking invention was awarded by Nobel prize in 2010.

Experimental studies of graphene are still restricted. Production of massive portions of the new substance still is a challenging synthetic problem. Many properties of graphene were *estimated*. Usually, there is not enough information for rigorous calculations, so we have to make assumptions and neglect unimportant factors. In this problem, you will estimate the adsorption properties of graphene.

1a. Estimate the specific surface of graphene open for adsorption in units m^2 /g . Consider that graphene plane is separated from any other solid or liquid substance.



The single layer of nitrogen molecules adsorbed on the outer surface of graphite is shown in Fig. 2. Assume that the same arrangement of nitrogen molecules is formed on a graphene surface.



Fig. 2. Nitrogen molecules N2 (grey circles) on the outer surface of graphite

1b. How many grams of nitrogen can be adsorbed on 1 gram of graphene assuming that the graphene layer is placed onto the surface of a solid support? Estimate the volume occupied by these nitrogen molecules after the complete desorption from 1 g of graphene (pressure 1 bar, temperature 298 K).

Calculations: $m_{N_2} = _____g$ $V_{N_2} = ______g$. Let us consider adsorption as a common chemical equilibrium

$$A_{gas} \rightleftharpoons A_{ads}, \tag{1}$$

(A_{gas} are molecules A in the gaseous state, A_{ads} are the same molecules on the surface) with the equilibrium constant *K*:

$$K = \frac{n_{A_{ads}} (\text{mol/m}^2)}{p_{A_{gas}} (\text{bar})}$$

(such assumption holds if a small number of molecules is adsorbed on the surface)

Adsorption properties of graphene can be estimated from the data for adsorption on a regular three-dimensional graphite. The enthalpy of adsorption (ΔH° of reaction (1)) of any molecule A on graphene is on average by 10% less negative compared to that on graphite. On graphite, the adsorbed molecule is bound more strongly due to the interaction with the lower graphene layers in the lattice (Fig. 1b) and hence the enthalpy of adsorption is more negative. The standard entropies of adsorption on graphene and graphite are assumed to be the same.

2. How many moles, *n*, of CCl₄ are adsorbed on 1 g of graphene at $p(\text{CCl}_4) = 10^{-4}$ bar if $2.0 \cdot 10^{-7}$ mol of CCl₄ are adsorbed on 1 m² of graphite at $p(\text{CCl}_4) = 6.6 \cdot 10^{-5}$ bar? Assume that graphene is placed onto the surface of a solid support and the interaction of CCl₄ with the support does not change the enthalpy of adsorption of CCl₄ on graphene. The temperature in both cases is 293 K. ΔH^0 of adsorption of CCl₄ on graphite is -35.1 kJ/mol.

Calculations: n(CCl₄) = _____

The graphene films are expected to be sensitive gas detectors. If 10^9 particles of a gas are adsorbed on 1 cm² of a graphene surface this is enough to measure an electrical resistivity change of the graphene layer and to detect the presence of a gas in the environment.

3. Determine the minimal content of ethane, C_2H_6 , in the air (in mol.%) at atmospheric pressure (T = 293K) at which a graphene sensor will detect this gas. The known data for the adsorption of alkanes on graphite are shown in Fig 3. Assume that air doesn't affect the adsorption properties of ethane.



Fig. 3. Thermodynamic properties for adsorption of alkanes on a graphite surface. (a) $\ln K \{ \text{mol/m}^2/\text{bar} \}$ as a function of $\ln M (M - \text{molecular mass of alkane in g/mol})$; (b) ΔH° of adsorption as a function of $\ln M$. Linear dependences are assumed in both cases

Calculations:

Content of $C_2H_6 =$ _____ mol.%

Problem 6. Cyclopropanes. So simple. So fancy... (8 points)

Question	1	2	3	Total
Marks	8	22	70	100

Cyclopropanes bearing donor and acceptor substituents at the neighboring C-atoms, for example, **A**, demonstrate high reactivity behaving similar to 1,3-zwitterion **B**.



Thus, A1 (X = 4-OMe) undergoes the three-membered ring opening in the Lewis acid-catalyzed reaction with 1,3-dimethoxybenzene as a nucleophile giving the product C.

1. Write down structural formula of C.

Structural formula of C:					

A1 participates in cycloadditions, annulations, oligomerizations, and other processes. Thus, formal [3+2]-cycloaddition between A1 and 4-methoxybenzaldehyde leads to a five-membered ring in **D**. Decarboxylation of all carboxylic groups in **D** produces **E** (C₁₈H₂₀O₃), the molecule of the latter having a plane of symmetry.



2. Write down structural formulae of **D** and **E** indicating their stereochemistry.

D	E

Also, **A** can undergo various transformations in the absence of any reaction partners except catalysts. Some transformations typical of **A1** are shown in the Scheme below.



To determine the structures of **F-J**, a set of physico-chemical data was obtained (see Table 1 for some results). It was found that:

a) **F** and **G** have the same molecular formula as **A1**;

b) G is formed as the most stable stereoisomer;

c) **H** and **I** are structural isomers;

d) **H** is formed as a single diastereomer with C_2 axis of symmetry (the molecule looks the same after rotation through the angle of 180°);

e) I is formed as a mixture of two diastereomers;

f) \mathbf{J} is naphthalene derivative.

In the process leading to **I**, one molecule of **A1** demonstrates the described above common reactivity (analogous to that of **B**). The other molecule of **A1** behaves differently. Also, the latter behavior is demonstrated by cyclopropane **A2** (dimethyl 2-(3,4,5-trimethoxyphenyl)cylopropane-1,1-dicarboxylate; X in $\mathbf{A} = 3,4,5$ -(MeO)₃) when treated with SnCl₄ affording **K** as a mixture of two diastereomers. The major isomer has the center of symmetry. Similar reactivity is shown by **A2** in Sn(OTf)₂-catalyzed reaction with **G** furnishing **L**.

$$\mathbf{K} \stackrel{\mathrm{SnCl}_4}{\longleftarrow} \mathbf{A2} \stackrel{\mathbf{G}}{\xrightarrow{}} \mathbf{I}$$

Table 1. Information about the compounds.

	Ratio	of the 1	groups	Empirical formula		
		١	Non-aromatic	Aromatic		
	CH	CH ₂	CH ₃	OH	CH	
A1	1	1	1+1+1	0	2+2	$(C_{14}H_{16}O_5)_n$
F	1	1	1+1+1	0	2+2	$(C_{14}H_{16}O_5)_n$
G	1+1+1	0	2+1	0	2+2	$(C_{14}H_{16}O_5)_n$
Η	1	1	1+1+1	0	2+2	$(C_{14}H_{16}O_5)_n$
Ι	1+1+1	1+1	2+1+1+1+1	0	2+2+1+1+1	$(C_{14}H_{16}O_5)_n$
J	0	0	1+1	1	1 + 1 + 1 + 1 + 1	$(C_{13}H_{12}O_4)_n$
K	1+1	1	2+1+1+1	0	1	$(C_{16}H_{20}O_7)_n$
L	1+1+1+1+1	1	2+2+1+1+1+1	0	2+2+1	$(C_5H_6O_2)_n$

3. Write down the structural formulae of **F-J**, **L** and the major isomer of **K**.

F	G
ľ	0
H	I
	-
Т	K (major isomer)
J	K (major isomer)
J	K (major isomer)
J	K (major isomer)
J	K (major isomer)
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J	K (major isomer)
J	K (major isomer)
J	K (major isomer)
J	K (major isomer)
L	K (major isomer)

Problem 7. Diverse permanganatometry (8 points)

Quest.	1	2	3			4		5	Total	
			a	b	c	d		a	b	
Marks	2	2	4	2	2	6	7	7	2	34

The amount of many reducing agents can be determined by permanganatometric titration in alkaline medium allowing permanganate ion reduction to manganate.

1. Write down the ionic equation of the reaction for formate titration with permanganate in an aqueous solution containing ~0.5 M NaOH.

Titration with permanganate in alkaline medium is often supplemented by addition of a barium salt, which leads to precipitation of manganate as $BaMnO_4$.

2. Which side redox processes involving manganate is suppressed by the barium salt? Write down an example of equation of the corresponding reaction.

10.00 mL (V_{Mn}) of 0.0400 M (c_{Mn}) KMnO₄ solution was placed in each of flasks **A**, **B**, and **C** and different reactions were conducted in each flask.

3. To flask **A**, a sample solution containing unknown amount of crotonic acid (CA) CH_3 -CH=CH–COOH (m_{CA}), an alkali and barium nitrate (both in an excess) were added, and the reaction mixture was incubated for 45 min. It is known that each molecule of crotonic acid loses 10 electrons under the experiment conditions. The molar mass of CA is 86.09 g/mol.

a) Write down the total ionic equation of the reaction.

8.00 mL (V_{CN}) of 0.0100 M (c_{CN}) potassium cyanide solution was further added to the incubated mixture. This resulted in completion of the following reaction:

$$2Ba^{2+} + 2MnO_4^{-} + CN^{-} + 2OH^{-} = 2BaMnO_4 + CNO^{-} + H_2O$$

BaMnO₄ precipitate was then filtered off, and the excess of cyanide in the filtrate was titrated with 0.0050 M (c_{Ag}) AgNO₃ solution till detectable precipitation was observed. Note that both CN⁻ and CNO⁻ are analogs of halide ions, but CNO⁻ affords soluble silver salt.

b) Give the formula for the complex formed when Ag^+ ions were initially added to the cyanide solution (until the precipitate was formed).

c) Give the formula of the precipitate formed.

d) Calculate the mass of crotonic acid (in mg) if 5.40 mL (V_{Ag}) of the silver salt solution was consumed for the titration to the endpoint.

4. Another sample with different concentration of crotonic acid and alkali (in an excess) were added to flask **B**, this mixture lacking barium salt. An excess of KI (instead of cyanide) was added as a reducing agent. The mixture was further acidified, and the iodine evolved was titrated with 0.1000 M (c_s) thiosulfate solution. 4.90 mL (V_{s1}) of the titrant was used to reach the endpoint.

Calculate the mass of crotonic acid (in mg).

5. A sample containing tin(II) was added to flask C, and the medium was adjusted to weak alkaline. Tin(II) was quantitatively oxidized to $Sn(OH)_6^{2-}$, whereas a precipitate formed as a result of permanganate reduction. The precipitate was isolated, washed off, dried at 250°C, weighed (the mass of the water-free precipitate (m_{prec}), representing a binary compound Mn_xO_y , was of 28.6 mg), and dissolved in H₂SO₄ in the presence of an excess of potassium iodide. The evolved iodine was titrated with 0.1000 M thiosulfate solution. 2.50 mL (V_{S2}) of the latter was consumed to attain the endpoint.

a) Determine x and y. Write down the reaction of precipitation.

b) Calculate the mass of tin in the sample (in mg).

Problem 8. Unique life of archaea (8 points)

Question	1	2	3	4	5	6	7	8	9		Total
									а	b	
Marks	2	7	3	8	4	4	5	4	3	5	45

Archaea (or archaebacteria) are single-celled microorganisms significantly differing from bacteria and eukaryotes at the molecular level.

Enzymatic reaction of methylamine with water is the major energy source for some archaea. In a particular experiment, an archaea strain was cultivated at pH 7 under anaerobic (oxygen free) conditions with the nutrient medium containing ¹³CH₃NH₂ as the only energy source. After a certain incubation period, the gas over the archaea culture was sampled and analyzed. It was found that the gas contains two substances **A** and **B** in the molar ratio of 1.00:3.00 correspondingly. The sample density rel. H₂ is of 12.0.

1. Calculate the volume fractions (in %) of **A** and **B** in the mixture.

2. Determine **A** and **B** if there is no N atoms in gas collected.

Your work:

Α	В

3. Write down the equation of enzymatic reaction of methylamine with water described in the above experiment using predominant form of each species.

Enzymes containing the residue of α -amino acid **X** are found in many archaea. It is known that **X**:

- is composed of atoms of 4 elements;
- is 18.8 % oxygen by mass;
- possesses the single individual tRNA and is incorporated into proteins during translation.

Amino acid *L*-lysine (see the structure in scheme below) was identified as the **X** precursor in archaea. All C and N atoms found in **X** originate from two starting lysine molecules. Different isotope-labeled *L*-lysines were introduced into a model system to clarify the biosynthetic pathways of **X**. The results are summarized in the table.

Isotope composition of <i>L</i> -lysine	Molecular mass (rounded to integer) of the X residue [RCH(NH ₂)CO], bound to tRNA,				
	g/mol				
Normal	238				
All carbons ¹³ C, all nitrogens ¹⁵ N	253				
ε-Amino group with ¹⁵ N	239				

4. Determine the molecular formula of **X**.

scheme equal 1.



X is biosynthesized in archaea according to the following scheme (E_1 – E_3 – enzymes):



At the first step, lysine is transformed into its structural isomer (α -amino acid, **C**), whereas **D** contains a peptide bond, and **E** a formyl group [$-C_{H}^{O}$]. All reaction coefficients in the above

5. Give the chemical formula of **C**, **D** and **E**. From the reaction types given hereunder, choose (tick) **only one** corresponding to the E_3 catalyzed reaction.

Calculations:		
С	D	E
□ Oxidative deamination;		
□ Decarboxylation;		
□ Intermolecular deamination	;	
\Box Hydroxylation;		
Peptide bond hydrolysis.		

X contains the fragment:



R is a massive substituent (M>100 g/mol). The 3^{rd} C atom is non-asymmetric, 4^{th} and 5^{th} C atoms are stereogenic centers. All C atoms in the cycle are bound with at least one H atom. Each substituent (H, Me and R) is found only once.

6. Determine the positions of substituents H, Me, and R.

Your work:

7. Draw structural formulae of C and X with stereochemical details. There are no stereo centers affected on the way from C to X. Mark every stereocenter of X with either R or S.



Only one codon is responsible for incorporation of \mathbf{X} residues into proteins in archaea. The nitrogen bases forming this codon contain two exocyclic amino groups and three exocyclic oxygen atoms in total.



8. Fill in the hereunder table to determine the nucleotide composition of the codon encoding **X**. **Tick only one box in each line**.

Your work:					
Nitrogen		The num	ber of bases in	the codon	
base	1	2	3	0 or 1	1 or 2
А					
С					
G					
U					

The fragment of mRNA coding sequence given below contains the codons encoding \mathbf{X} residue incorporation into an archaea enzyme:

5'...AAUAGAAUUAGCGGAACAGAGGGUGAC...3'

9a. Using the table of the genetic code, decide how many amino acid residues are incorporated into the enzyme chain due to this fragment translation.

Your v	work:
--------	-------

Number of amino acids =

9b. Write down the amino acid sequence translated from this fragment. Note that more than one X residue is found in the fragment.

Fill in the boxes with the amino acid abbreviations (from N- to C-terminus).

Note that the number of boxes is excessive. If there is more than one possibility, write all separated by "/". If the translation is stopped in a particular position, write "STOP" and leave all the boxes to the right empty.

Your	work:						
		1	 	 1			1
							_

(a) RNA Codons for the Twenty Amino Acids

		second	Uase			-
	U	С	Α	G		
	Phe	Ser	Tyr	Cys	U	
IJ	Phe	Ser	Tyr	Cys	С	
U	Leu	Ser	STOP	STOP	Α	
	Leu	Ser	STOP	Trp	G	
	Leu	Pro	His	Arg	U	
C	Leu	Pro	His	Arg	С	
U	Leu	Pro	Gln	Arg	Α	T
	Leu	Pro	Gln	Arg	G	nro
	Ile	Thr	Asn	Ser	U	l ba
	Ile	Thr	Asn	Ser	С	se
A	Ile	Thr	Lys	Arg	Α	
	Met(start)	Thr	Lys	Arg	G	
	Val	Ala	Asp	Gly	U	
C	Val	Ala	Asp	Gly	С	
G	Val	Ala	Glu	Gly	Α	
	Val	Ala	Glu	Gly	G]

second base

Amino acid abbreviations:

Ala = Alanine Arg = Arginine Asn = Asparagine Asp = Aspartic acid Cys = Cysteine Glu = Glutamic acid Gln = Glutamine Gly = Glycine His = Histidine Ile = Isoleucine Leu = Leucine Lys = Lysine Met = Methionine Phe = Phenylalanine Pro = Proline Ser = Serine Thr = Threonine Trp = TryptophanTyr = Tyrosine Val = Valine

Group	1	2		3	4	5	6	7	8	9	10	11	12	13	14	15	16	17	18
Period																			
	1																		2
1	Н																		He
	1.008		i.																4.0026
0	3	4												5	6	7	8	9	10
2	LI	De												D	12.011	14.007	15.000	19.009	1NC
	0.94	9.0122												10.81	12.011	14.007	16	17	20.180
3	Na	Ma												ΔΙ	Si	P	S	č	Δr
	22.990	24.305												26.982	28.085	30.974	32.06	35.45	39.948
	19	20		21	22	23	24	25	26	27	28	29	30	31	32	33	34	35	36
4	K	Ca		Sc	Ti	V	Cr	Mn	Fe	Co	Ni	Cu	Zn	Ga	Ge	As	Se	Br	Kr
	39.098	40.078		44.956	47.867	50.942	51.996	54.938	55.845	58.933	58.693	63.546	65.38	69.723	72.63	74.922	78.96	79.904	83.798
	37	38		39	40	41	42	43	44	45	46	47	48	49	50	51	52	53	54
5	Rb	Sr		Y	Zr	Nb	Mo	Тс	Ru	Rh	Pd	Ag	Cd	In	Sn	Sb	Те	I	Xe
	85.468	87.62		88.906	91.224	92.906	95.96	[97.91]	101.07	102.91	106.42	107.87	112.41	114.82	118.71	121.76	127.60	126.90	131.29
0	55	56		71	72	73	74	75	76	77	78	79	80	81	82	83	84	85	86
6	US	Ва		LU	HI	18	VV	Re	Us	Ir	Pt	Au	Hg	11	PD	BI	PO	At	Rn
	132.91	137.33		1/4.9/	1/8.49	180.95	183.84	180.21	190.23	192.22	195.08	196.97	200.59	204.38	207.2	208.98	[208.98]	[209.99]	[222.02]
7	67 Er	Da	**	103 r	104 Df	Dh	001 Sa	Bh	He	ТU9 МЛ+	De	Pa	Cn	I lut	FI	llun			
,	[223 [12]	1226 031		[262 11]	[265 12]	[268 13]	1271 131	12701	[277 15]	[276 15]	[281 16]	1280 161	[285 17]	[284 18]	[289 19]	[288 19]	[293]	[294]	[294]
	[[[[[[[]]]]]]]]	[220.00]		[202.11]	[200.12]	[200.10]	[271.10]	[2,0]	[[Elificity]	[[210.10]]	[201.10]	[200.10]	[200.17]	[204.10]	[200.10]	[200.10]	[200]	[204]	[204]
			i	-		-											-		
*Lanthanoida			*	57	58	59	60 Not	61 Dura	62 0.000	63	64 Cd	65 TL	66 Dv	67	68 E w	69 T.m.	70 Vh		
Lanunanoida		0.00	La	Ce	P1	114.04	Pm	50.00	EU	457.05	G I	Dy tea so	164.00	197.00	1 111	4 70 OF			
				130.91	140,12	Q1	02	[144.91] 02	150.30 QZ	05	137.23 QB	00.90	02.00	04.95 QQ	107.20	100.95	102		
**Actinoids			**	Ac	Th	Pa	U	Np	Pu	Am	Cm	Bk	Cf	Es	Fm	Md	No		
				[227.03]	232.04	231.04	238.03	[237.05]	[244.06]	[243.06]	[247.07]	[247.07]	[251.08]	[252.08]	[257.10]	[258.10]	[259.10]		